Applied Genome Research

De novo transcriptome assembly

205048 & 205049



Generates initial assembly of dominant isoforms

(https://github.com/trinityrnaseq/trinityrnaseq/wiki)

Constructs graph of common sequences and unique sequences of different isoforms

Resolves graph and reports separate isoforms (final assembly)

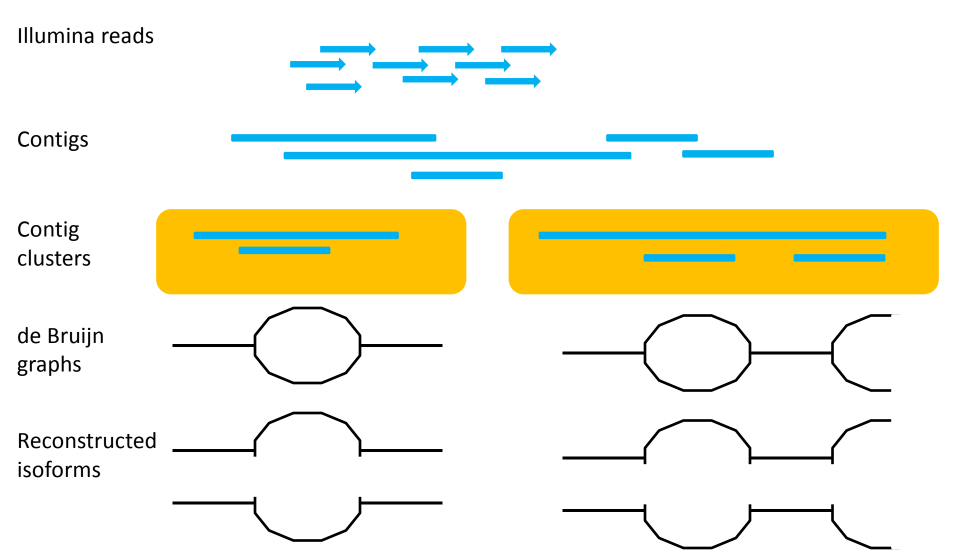
Running Trinity

```
$ Trinity \
--normalize_reads \
--seqType fq \
--max_memory 20G \
--single <INPUT>.fastq \
--CPU 6 \
--output <OUTPUT_DIRECTORY>
```

Trinity on cluster

```
#!/bin/bash
                         r/trinityrnaseq-Trinity-v2.4.0/Trinity \
echo "/c
--normalize reads \
--seqType fq \
--max memory 10G \
--left 1P.fastq,left 1P.fastq \
--right 2P.fastq,2P.fastq \
--CPU 4 --output <SOME DIRECTORY>" \
| qsub \
-cwd \
-N iGEM trin \
-l vf=10G -l arch=lx-amd64 -l idle=1 \
-P fair share \
-pe multislot 20 \
-o output.txt \
-e error.txt
```

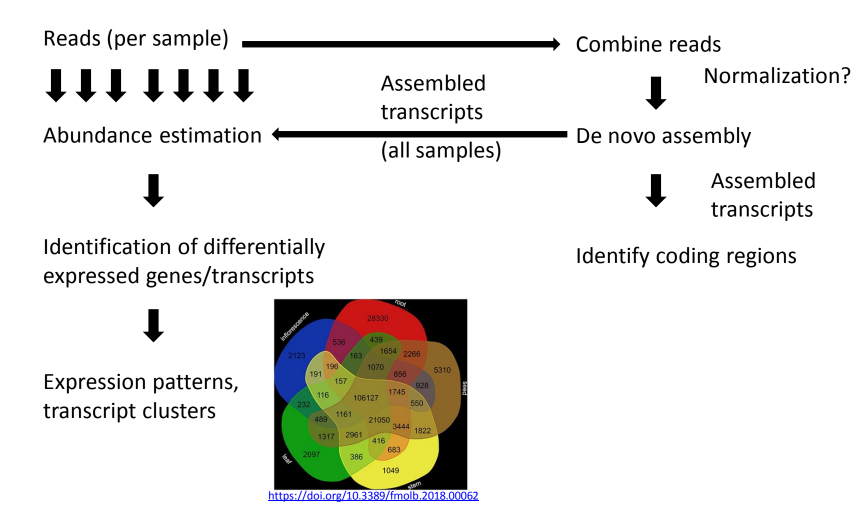
Components of Trinity



Boas Pucker

5

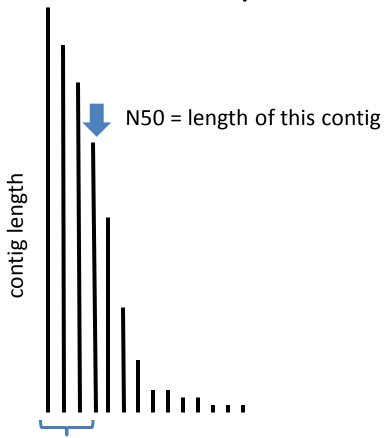
General analysis concept



Best practice workflow

- 1) Generate resource report:
 - trinityrnaseq-Trinity-v2.4.0/trinity-plugins/COLLECTL/examine_resource_usage_profiling.pl collect
- 2) Check assembly for full length transcripts => BLASTx vs. nr
- 3) Analyze tophit coverage
- 4) Check integrated hits via bowtie mapping against assembly
- 5) Calculate Nx stats e.g. Ex90N50 (N50 is not useful)
- 6) Run BUSCO to check assembly completeness
- 7) Run Interproscan to assign GO terms

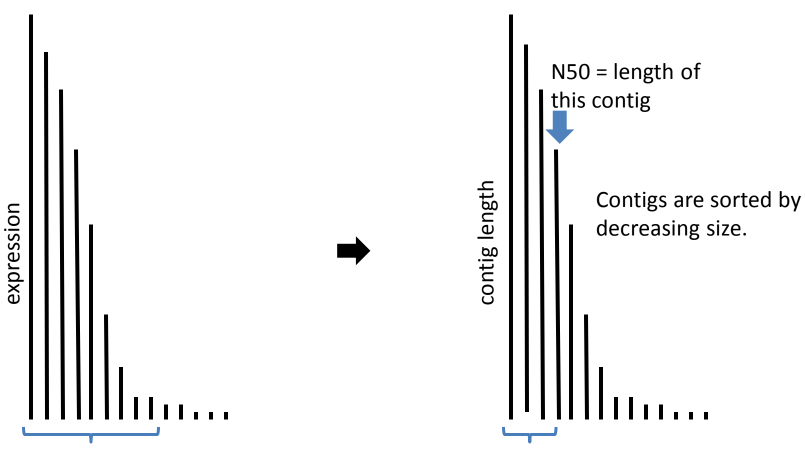
Assembly evaluation – Nx for continuity quantification



Contigs are sorted by decreasing size.

Contigs sum up to 50% of total assembly size

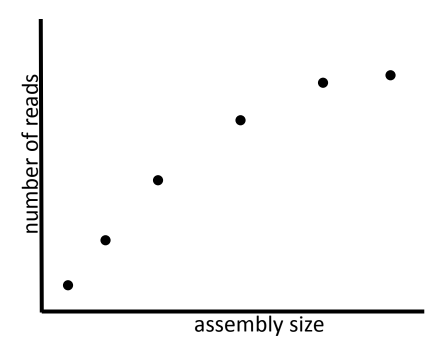
Assembly evaluation – Ex90N50



Sorting contigs by expression and selecting all sequences that account for 90% of all expression

Contigs sum up to 50% of selected assembly fraction

Saturation of assembly size



BUSCO

- "quantitative measures for the assessment of genome assembly, gene set, and transcriptome completeness, based on evolutionarily-informed expectations of gene content from near-universal single-copy orthologs"
- https://busco.ezlab.org/
- Applications: assembly completeness assessment, estimation of heterozygosity, optimization of gene prediction, identification of paralogs, ...

Gene Ontology (GO) terms

- http://geneontology.org/
- Computational representation of biological knowledge
- Over 40k biological concepts
- Used for automatic annotation of predicted genes
- GO enrichment analyses (e.g. in RNA-Seq studies)

Construct GFF3

- One feature/entry in GFF3 file is created per sequence in FASTA file
- Length of sequences is used to determine start/end
- Running number is used to generate unique IDs

EXERCISE

- 1) Run 'contig_stats.py' on Trinity.fasta!
- 2) Use STAR to map all WT and 3xmyb reads to assembly!
- 3) Construct reference file for read counting via 'fasta2gff.py'!
- 4) Use featureCounts to get expression values!
- 5) Construct heatmap via 'construct_heatmap.py'! (see tipps)

Construct Heatmap

Add your own file paths and file names!

Run script like always: python construct_heatmap.py